## Copyright (c) 1993 - 2004 GenCore version 5.1.6 Compugen Ltd.

OM nucleic - nucleic search, using sw model

Run on: August 2, 2004, 14:00:47; Search time 32049 Seconds

(without alignments)

11691.489 Million cell updates/sec

Title: US-10-038-854-37

Perfect score:

Sequence: 1 tttggcctcgggccagaatt.....actgtatttaactaacttta 8645

Scoring table: IDENTITY\_NUC

Gapop 10.0 , Gapext 1.0

Searched: 3470272 seqs, 21671516995 residues

Total number of hits satisfying chosen parameters: 6940544

Minimum DB seq length: 0
Maximum DB seq length: 200000000

Post-processing: Minimum Match 0% Maximum Match 100%

Listing first 45 summaries

Database GenEmbl: \*

6 :: : gb\_ba:\*
gb\_htg:\*
gb\_in:\*
gb\_om:\*
gb\_ov:\*
gb\_pat:\*

7 :

gb\_ph:\*
gb\_p1:\*
gb\_pr:\*

gb\_ro:\*

```
us-10-038-854-37
```

```
40:
               39:
                    38:
                           37:
                                 36:
                                        35:
                                               33
4:
                                                            32:
                                                                  31:
                                                                         29:
30:
                                                                                      28:
                                                                                             24:
25:
26:
                                                                                                                      23:
                                                                                                                             22:
                                                                                                                                          20:
                                                                                                                                                       18:
                                                                                                                                                                           15:
                                                                                                                                                                                  14:
                                                                                                                                                                                        13:
                                                                                                                                   21:
                                                                                                                                                 19:
                                                                                                                                                              17:
                                                                                                                                                                     16:
                                                                                                                                                                                               12:
                                                                 em_vi:*
em_htg_hum:*
em_htg_inv:*
                                      em_htg_mus:*
em_htg_pln:*
em_htg_rod:*
                                                                                                                                                                   gb_sts:*
gb_un:*
gb_vi:*
em_ba:*
em_fun:*
                                                                                                                                               em_in:*
em_mu:*
em_htgo_mus:*
em_htgo_other:*
            em_htgo_hum: *
                                                          em_htg_other:*
                                                                                    em_un:*
                                                                                           em_sts:*
                                                                                                 em_ro:*
                                                                                                        em_p1:*
                                                                                                              em_ph:*
                                                                                                                     em_pat:*
                                                                                                                            em_ov:*
                                                                                                                                  em_or:*
                                                                                                                                         em_om:*
                                                                                                                                                            em_hum:*
                   em_sy:*
                          em_htg_vrt:*
                                em_htg_mam:*
```

score greater than or equal to the score of the result being printed, Pred. No. is the number of results predicted by chance to have a and is derived by analysis of the total score distribution.

#### SUMMARIES

Result Query
No. Score Match Length DB ID

Description

Page 2

50065 Sequer 76360 Sequer	$\circ$	6560 2157	30.1 24.9	2599 2155.4	3 3 9
00772 Homo	AF100772	8297	•	2854.8	37
48178 Homo	HSM808	7713	•	9	36
025410 Mus mu	0 AB02541	8373	•	99	<b>3</b> 5
38613 Gallus	GGA238	8118	•	3012.6	34
6980 Danic	AB0269	9264	35.5	9	33
953 Homo	AB0329	7781	•	3158	32
5869 Homo	AK125	3394	•	3235.6	31
Mus	0 AI	9722	•	2	30
748	AK00	3270	37.8	3270	29
Prime	BDJ	3270	•	3270	28
Sequenc	AX8	3270	٠	3270	27
AX600210 Sequence	⊳	8645	39.8	44	26
025413		8585	•	4.5	25
		9729	40.0	3453.8	24
		9826	•	48	23
AX556500 Sequence		8354	40.5	5	22
086607		6898	•	ភ	21
ťΩ		8689	•	ក្ន	20
025411		8797	•	3545.6	19
50066		8797	•	2.0	18
0063		8797	•	5.	17
50068 Sequenc		40	•	56	16
031		8409	•	5	15
51 Sequ	6 AX675551	8438	41.5	5	14
1336		3614	41.8	3614	13
75 Prim		3614	•	3614	12
5 Sequ	AX87652	3614	•	3614	11
921803 Sequer	AX92180	8575	43.0	71	10
9	AB02697	8816	•	2	9
22513 Mus	AK1225	5804	•	4561.4	œ
1 8880	AB0408	5309	•	305.	7
ū	AF1954	7816	•	749.	σ
5412	AB025	9	•	789.	σ
359	AX6623	48	•	95.	4
57 S	AX66235	8473	N	000.	ω
353 S	AX6623	σ	99.7	8616.6	N
623	AX66235	64	100.0	64	1
	1 1 1 1 1 1 1	١			

)
)
1
ř
Ø
- 1
$\vdash$
0
- 1
0
ω
ω
1
ω
ഗ
4
- 1
ω
J
-

45 1614.8 18.7 184032 9 AC079226 AC	19.2 5583 10 AK122490	1671.2 19.3 8624 9 AB037723	1978.8 22.9 8993 9 HSM806812	9 AK027473	2155.4 24.9 2157 6 BD156088
AC079226 Homo sap	AK122490 Mus	AB037723 Homo sap	BX640737 Homo	AK027473 Homo	BD156088 Primer f

fo api api api api

Search completed: August Job time : 32211 secs 3, 2004, 06:21:49

Copyright (c) 1993 - 2004 GenCore version 5.1.6 Compugen Ltd

OM nucleic - nucleic search, using sw model

Run on: August 2, 2004, 20:38:12; Search time 3396 Seconds

12481.639 Million cell updates/sec (without alignments)

Title: US-10-038-854-37

Perfect score: 8645

Sequence: 1 tttggcctcgggccagaatt.. ..actgtatttaactaacttta 8645

Scoring table: IDENTITY\_NUC

Gapop 10.0 , Gapext 1.0

Searched: 3222919 seqs, 2451570024 residues

Total number of hits satisfying chosen parameters: 6445838

Minimum DB seq length: 0
Maximum DB seq length: 200000000

Post-processing: Minimum Match 0%

Listing first 45 summaries

Maximum Match 100%

Database

Published\_Applications\_NA:\*
/cgn2\_6/ptodata/2/pubpna/US07\_PUBCOMB.seq:\*

5:

7 : 9 /cgn2\_6/ptodata/2/pubpna/PCT\_NEW\_PUB.seq:\*/cgn2\_6/ptodata/2/pubpna/US06\_NEW\_PUB.seq:\*/cgn2\_6/ptodata/2/pubpna/US06\_PUBCOMB.seq:\*/cgn2\_6/ptodata/2/pubpna/US07\_NEW\_PUB.seq:\*/cgn2\_6/ptodata/2/pubpna/PCTUS\_PUBCOMB.seq:\*/ cgn2\_6/ptodata/2/pubpna/US08\_NEW\_PUB.seq:\*

œ .. cgn2\_6/ptodata/2/pubpna/US08\_PUBCOMB.seq:\*

10: cgn2\_6/ptodata/2/pubpna/US09A\_PUBCOMB.seq:\*

cgn2\_6/ptodata/2/pubpna/US09B\_PUBCOMB.seq:\*

# us-10-038-854-37

13

19:	18:	17:	16:	15:	14:	13:	12:	11:
6/ptodata/2/pubpna/US60_PUB	<pre>/cgn2_6/ptodata/2/pubpna/US60_NEW_PUB.seq:*</pre>	_6/ptodata/2/pubpna/US10_	_6/ptodata/2/pubpna/US10C	_6/ptodata/2/pubpna/US10B	n2_6/ptodata/2/pubpna/US10A_PUBCOMB.s	6/ptodata/2/pubpna/US09_NEW	<pre>/cgn2_6/ptodata/2/pubpna/US09_NEW_PUB.seq:*</pre>	<pre>/cgn2_6/ptodata/2/pubpna/US09C_PUBCOMB.seq:*</pre>

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

### SUMMARIES

1	0	9	ω	7	Q	U		ω	N	1	0		80	7	6					٣	Result No.
3466.4	482.	482.	502.	502.	518.	518.	3545.6	545.	545.	545.	566.	6	3589.8	61	61	3714.8	895.	000.	8616.6	8645	Score
40.1	40.3	40.3	40.5	40.5	40.7	0	41.0	۳.	1.	1.	۲.	۳.	41.5	•	۲.	43.0	μ.	92.5	99.7	100.0	% Query Match
9	9826	Ν	5	8354	æ	8689	8797	8797	9	9	8409	8409	8438	$\vdash$	$\mathbf{L}$		α	8473	7	8645	Length
16	10	9	13	13	10	9	10	10	9	9	10	9	13	13	13	13	17	17	17	17	DB
US-10-144-194A-81	US-09-800-198-7	-808-60	US-10-029-020-13	-10-383	-09-800-198-	09-808-602-7	US-09-800-198-65	9-800-198-	US-09-808-602-77	9-808-6	50-	02-7	2-8	-118-17	2-887-1	2-	8-854-4			US-10-038-854-37	ID
Sequence 81, Appl	Sequence 7, Appli	Sequence 7, Appli	Sequence 13, Appl.	Sequence 43, Appl	Sequence 66, Appl	Sequence 78, Appl	Sequence 65, Appl	Sequence 62, Appl	Sequence 77, Appl	Sequence 74, Appl	Sequence 67, Appl	Sequence 79, Appl	Sequence 1, Appli	Sequence 1743, Ap	Sequence 1743, Ap	Sequence 143, App	Sequence 41, Appl	Sequence 39, Appl	Sequence 35, Appl	Sequence 37, Appl	Description

٠.		~	.0	•	_	v	~	~	٥,	٥,		~		•	_	•	~	~	٠,	٠,				
421.6	423	423	556.6	584.4	584.4	584.4	617.8	621.2	622	622	663.8	663.8	706	706	766.4	912.8	2599	2599	2854.8	3249.6	3264.8	3453.8	3453.8	
4.9	4.9	4.9	6.4	6.8	6.8	6.8	7.1	7.2	7.2	7.2	7.7	7.7	8.2	8.2	8.9	10.6	30.1	30.1	33.0	37.6	37.8	40.0	40.0	
452	829	829	1973	1755	1755	1755	1371	1476	1392	1392	1399	1399	2496	2496	791	3217	6560	6560	12880	8355	9058	9729	9729	
9	13	13	9	16	15	10	13	13	13	13	13	13	10	9	13	15	10	9	16	13	16	10	9	
US-09-563-817-960	US-10-383-201-61	US-10-383-201-47	US-09-864-761-4526	US-10-384-974-17	US-10-004-415-17	US-09-998-966-17	US-10-383-201-51	US-10-383-201-41	US-10-383-201-53	US-10-383-201-45	US-10-383-201-59	US-10-383-201-49	US-09-800-198-63	US-09-808-602-75	US-09-823-245A-85	US-10-198-846-13976	US-09-800-198-64	US-09-808-602-76	US-10-295-027-927	US-10-383-201-55	US-10-144-194A-79	US-09-800-198-12	US-09-808-602-12	us-10-038-854-37
Sequence 960, App	Sequence 61, Appl	Sequence 47, Appl	Sequence 4526, Ap	Sequence 17, Appl	Sequence 17, Appl	17,	51,	41,	Sequence 53, Appl	Sequence 45, Appl	Sequence 59, Appl	Sequence 49, Appl	Sequence 63, Appl	Sequence 75, Appl	Sequence 85, Appl	Sequence 13976, A	Sequence 64, Appl	76, Aj	Sequence 927, App	Sequence 55, Appl	Sequence 79, Appl	Sequence 12, Appl	Sequence 12, Appl	

a

### ALIGNMENTS

Ω

41

Search completed: August 3, 2004, 12:53:36 Job time : 3692 secs